

# An Analytical Study of a Structured Overlay in the presence of Dynamic Membership

Supriya Krishnamurthy<sup>1</sup>, Sameh El-Ansary<sup>1</sup>, Erik Aurell<sup>1,2</sup> and Seif Haridi<sup>1,3</sup>

<sup>1</sup> Swedish Institute of Computer Science (SICS), Sweden

<sup>2</sup> Department of Physics, KTH-Royal Institute of Technology, Sweden

<sup>3</sup> IMIT, KTH-Royal Institute of Technology, Sweden

{supriya,sameh,eaurell,seif}@sics.se

**Abstract**— In this paper, we present a complete analytical study of dynamic membership (aka churn) in structured peer-to-peer networks. We use a master-equation-based approach, which is used traditionally in non-equilibrium statistical mechanics to describe steady-state or transient phenomena. We demonstrate that this methodology is in fact also well suited to describing structured overlay networks by an application to the Chord system. For any rate of churn and stabilization rates, and any system size, we accurately account for the functional form of: the distribution of inter-node distances, the probability of network disconnection, the fraction of failed or incorrect successor and finger pointers and show how we can use these quantities to predict both the performance and consistency of lookups under churn. Additionally, we also discuss how churn may actually be of different ‘types’ and the implications this will have for structured overlays in general. All theoretical predictions match simulation results to a high extent. The analysis includes details that are applicable to a generic structured overlay deploying a ring as well as Chord-specific details that can act as guidelines for analyzing other systems.

## I. INTRODUCTION

An intrinsic property of Peer-to-Peer systems is the process of never-ceasing dynamic membership. Structured Peer-to-Peer Networks (aka Distributed Hash Tables (DHTs)) have the underlying principle of arranging nodes in an overlay graph of known topology and diameter. This knowledge results in the provision of performance guarantees. However, dynamic membership continuously “corrupts/churns” the overlay graph and every DHT strives to provide a technique to “correct/maintain” the graph in the face of this perturbation.

Both theoretical and empirical studies have been conducted to analyze the performance of DHTs undergoing “churn” and simultaneously performing “maintenance”. Liben-Nowell et. al [7] prove a lower bound on the maintenance rate required for a network to remain connected in the face of a given dynamic membership rate. Aspnes et. al [3] give upper and lower bounds on the number of messages needed to locate a node/data item in a DHT in the presence of node or link failures. The value of such theoretical studies is that they provide insights neutral to the details of any particular DHT. Empirical studies have also been conducted to complement these theoretical studies by showing how within the asymptotic bounds, the performance of a DHT may vary substantially

depending on different DHT designs and implementation decisions. Examples include the work of: Li et. al [6], Rhea et.al [9] and Rowstron et.al [4].

In this paper, we present a new approach to studying churn, based on working with master equations, a widely used tool wherever the mathematical theory of stochastic processes is applied to real-world phenomena [8]. We demonstrate the applicability of this approach to one specific DHT: Chord [10].

A master-equation description for a dynamically evolving system is achieved by first defining a *state* of the system. This is just a listing of the quantities one would need to know for the fullest description of the system. For Chord, the *state* could be defined as a listing of how many nodes there are in the system and what the state (whether correct, incorrect or failed) of each of the pointers of those nodes is. This information is not enough to draw a unique graph of network-connections (because for example, if we know that a given node has an ‘incorrect’ successor pointer, this still does not tell us which node it is pointing to). However, as we will see, beginning at this level of description is sufficient to keep track of most of the details of the Chord protocols.

Having defined a state, the master-equation is simply an equation for the evolution of the probability of finding the system in this state, given the details of the dynamics. The specific nature of the dynamics plays a role in evaluating all the terms leading to the gain or loss of this probability, *i.e.* keeping track of the contribution of all the events which can bring about changes in the probability in a micro-instant of time.

Using this formalism our results are accurate functional forms of the following: (i) The distribution of inter-node distances when the system is in equilibrium. This distribution is independent of any details of Chord and are applicable to any DHT deploying a ring. (ii) Chord-specific inter-node distribution properties. (iii) For every outgoing pointer of a Chord node, we systematically compute the probability that it is in any one of its possible states. This probability is different for each of the successor and finger pointers. We then use this information to predict other quantities such as (iv) the probability that the network gets disconnected, (v) lookup consistency (number of failed lookups), and (vi) lookup performance (latency). All quantities are computed as a function of the parameters involved and all results are verified

by simulations.

## II. RELATED WORK

Closest in spirit to our work is the informal derivation in the original Chord paper [10] of the average number of timeouts encountered by a lookup. This quantity was approximated there by the product of the average number of fingers used in a lookup times the probability that a given finger points to a departed node. Our methodology not only allows us to derive the latter quantity rigorously but also demonstrates how this probability depends on which finger (or successor) is involved. Further we are able to derive a precise relation relating this probability to lookup performance and consistency accurately at any value of the system parameters.

In the works of Aberer et.al [1] and Wang et.al [11], DHTs are analyzed under churn and the results are compared with simulations. However, the main parameter of the analysis is the probability that a random selected entry of a routing table is stale. In our analysis, we determine this quantity from system details and churn rates.

A brief announcement of the results presented in this paper, has appeared earlier in [5].

## III. OUR IMPLEMENTATION OF CHORD

**The Chord Ring.** The general philosophy of DHTs is to map a set of data items onto a set of nodes where the insertion and lookup of items is done using unique keys of items. Chord's realization of that philosophy is as follows. Peers and data items are given unique keys (usually obtained by a cryptographic hash of unique attribute like the IP address or public key for nodes, and filename or checksum for items) drawn from a circular key space of size  $\mathcal{K}$ . The Chord system dictates that the right place for storing an item is at the first alive node whose key succeeds the key of the item. Since we refer to nodes and items by their keys, in that way, the insertion and lookup of items becomes a matter of locating the right "successor" of a key. All nodes have successor and predecessor pointers. For  $N$  nodes, using only the successor pointers to lookup items requires  $\frac{1}{2}N$  hops on average.

**Fingers.** To reduce the average lookup path length, nodes keep  $\mathcal{M} = \log_2 \mathcal{K}$  pointers known as the "fingers". Using these fingers, a node can retrieve any key in  $O(\log N)$  hops. The fingers of a node  $n$  (where  $n \in 0 \dots \mathcal{K} - 1$ ) point to exponentially increasing distances of keys away from  $n$ . That is,  $\forall i \in 1..\mathcal{M}$ ,  $n$  points to a node whose key is equal  $n + 2^{i-1}$ . We denote that key by  $n.\text{fin}_i.\text{start}$ . However, for a certain  $i$ , there might not be a node in the network whose key is equal to  $n + 2^{i-1}$ . Therefore,  $n$  points to the successor of  $n + 2^{i-1}$  which we denote by  $n.\text{fin}_i.\text{node}$ .

**The Successor List** Moreover, each node keeps a list of the  $\mathcal{S} = O(\log(N))$  immediate successors as backups to its first successor we use the notation  $n.s$  to refer to this list and  $n.s_i$  to refer to the  $i^{\text{th}}$  element in the list. Finally we use the notation  $n.p$  to refer to the predecessor

**Stabilization, Churn & Steady State.** To keep the pointers up-to-date in the presence of churn, each node performs periodic stabilization of its successors and fingers. In our

analysis, we define  $\lambda_j$  as the rate of joins per node,  $\lambda_f$  the rate of failures per node and  $\lambda_s$  the rate of stabilizations per node. We carry out our analysis for the general case when the rate of doing successor stabilizations  $\alpha\lambda_s$ , is not necessarily the same as the rate at which finger stabilizations  $(1 - \alpha)\lambda_s$  are performed. In all that follows, we impose the steady state condition  $\lambda_j = \lambda_f$  unless otherwise stated. Further it is useful to define  $r \equiv \frac{\lambda_s}{\lambda_f}$  which is the relevant ratio on which all the quantities we are interested in will depend, e.g,  $r = 50$  means that a join/fail event takes place every half an hour for a stabilization which takes place once every 36 seconds. Throughout the paper we will use the terms  $\lambda_j \Delta t$ ,  $\lambda_f \Delta t$ ,  $\alpha\lambda_s \Delta t$  and  $(1 - \alpha)\lambda_s \Delta t$  to denote the respective probabilities that a join, failure, a successor stabilization, or a finger stabilization take place during a micro period of time of length  $\Delta t$ .

**Parameters.** The parameters of the problem are hence:  $\mathcal{K}$ ,  $N$ ,  $\alpha$  and  $r$ . All relevant measurable quantities should be entirely expressible in terms of these parameters.

**Simulation** Since we are collecting statistics like the probability of a particular finger pointer to be wrong, we need to repeat each experiment 100 times before obtaining well-averaged results. The total simulation sequential real time for obtaining the results of this paper was about 1800 hours that was parallelized on a cluster of 14 nodes where we had  $N = 1000$ ,  $\mathcal{K} = 2^{20}$ ,  $\mathcal{S} = 6$ ,  $200 \leq r \leq 2000$  and  $0.25 \leq \alpha \leq 0.75$ .

While the main outlines of the chord protocol are provided by its authors in [10], an exact analysis necessitates the provision of a deeper level of detail and adopted assumptions which we provide in the following subsections.

### A. Joins, Failures & Ring Stabilization

**Initialization.** Initially, a node knows its key and at least one node with key  $c$  that already exists in the network and is alive. The knowledge of such a node is assumed to be acquired through some out-of-band method. The predecessor  $p$ , successors ( $s_1..s$ ) and fingers ( $\text{fin}_{1..\mathcal{M}}.\text{node}$ ) are all assigned to *nil*.

**Joins** (Fig. 1). A new node  $n$  joins by looking up its successor using the initial random contact node  $c$ . It also starts its first stabilization of the successors and initializes its fingers.

**Stabilization of Successors** (Fig. 1). The function *fixSuccessors* is triggered periodically with rate  $\alpha\lambda_s$ . A node  $n$  tells its first alive successor  $y$  that it believes itself to be  $y$ 's predecessor and expects as an answer  $y$ 's predecessor  $y.p$  and successors  $y.s$ . The response of  $y$  can lead to three actions: *Case A.* Some node exists between  $n$  and  $y$  (i.e.  $n$ 's belief is wrong), so  $n$  prepends  $y.p$  to its successor list as a first successor and retries *fixSuccessors*.

*Case B.*  $y$  confirms  $n$ 's belief and informs  $n$  of  $y$ 's old predecessor  $y.p$ . Therefore  $n$  considers  $y.p$  as an alternative/initial predecessor for  $n$ . Finally,  $n$  reconciles its successor list with  $y.s$ .

*Case C.*  $y$  agrees that  $n$  is its predecessor and the only task of  $n$  is to update its successor list by reconciling it with  $y.s$ .

By calling *iThinkIamYourPred* (Fig. 1), some node  $x$  informs  $n$  that it believes itself to be  $n$ 's predecessor. If  $n$ 's

```

n.join(c)
  s1 = c.findSuccessor(n)
  fixSuccessors()
  initFingers(s1)
n.fixSuccessors()
  y = firstAliveSuccessor()
  {y.p, y.s} = y.iThinkIamYourPred(n)
  if (y.p ∈ ]me, y]) //Case A
    prepend(y.p)
    fixSuccessors()
  elseif (y.p ∈ ]y, me]) //Case B
    considerANewPred(y.p)
    reconcile(y.s)
  else //Case C: y.p == me
    reconcile(y.s)
n.firstAliveSuccessor()
  while (true)
    if (s1 == nil)
      //Broken Ring!!
    if (isAlive(s1))
      return (s1)
    ∀ i ∈ 1..(S - 1)
      si = si+1
    sS = nil
n.considerANewPred(x)
  if (isNotAlive(p)
    or (p == nil)
    or (x ∈ ]p, n])
    p = x
n.iThinkIamYourPred(x)
  if ((isNotAlive(p) or (p == nil))
    p = x
    return({s, x})
  if (x ∈ ]p, me])
    oldp = p
    p = x
    return({s, oldp})
  else
    return({s, p})
n.reconcile(s')
  for i = 1..(S - 1)
    si+1 = s'i
n.prepend(y)
  for i = S..2
    si = si-1
    s1 = y

```

Fig. 1. Joins and Ring Stabilization Algorithms

predecessor  $p$  is not alive or  $nil$ , then  $n$  accepts  $x$  as a predecessor and informs  $x$  about this agreement by returning  $x$ . Alternatively, if  $n$ 's predecessor  $p$  is alive (discovering that will be explained shortly in section III-C), then there are two possibilities: The first is that  $x$  is in the region between  $n$  and its current predecessor  $p$  therefore  $n$  should accept  $x$  as a new predecessor and inform  $x$  about its old predecessor. The second is that  $p$  is already pointing to  $x$  so the state is correct at both parties and  $n$  confirms that to  $x$  by informing it that  $x$  is the predecessor of  $n$ . In all cases the function returns a predecessor and a successors list.

The function *firstAliveSuccessor* (Fig. 1) iterates through the successors list. In each iteration, if the first successor  $s_1$  is alive, it is returned. Otherwise, the dead successor is dropped from the list and  $nil$  is appended to the end of the list. If the first successor is  $nil$  this means that all immediate successors are dead and that the ring is disconnected.

### B. Lookups and Stabilization of Fingers

**Stabilization of Fingers** (Fig. 2). Stabilization of fingers occurs at a rate  $(1 - \alpha)\lambda_s$ . Each time the *fixFingers* function is triggered, a random finger  $fin_i$  is chosen and a lookup for  $fin_i.start$  is performed and the result is used to update  $fin_i.node$ .

**Initialization of Fingers** (Fig. 2). After having initialized its first successor  $s_1$ , a node  $n$  sets all fingers with starts between  $n$  and  $s_1$  to  $s_1$ . The rest of the fingers are initialized by taking a copy of the finger table of  $s_1$  and finding an approximate successor to every finger from that finger table.

**Lookups** (Fig. 3). A lookup operation is a fundamental operation that is used to find the successor of a key. It is used by many other routines and its performance and consistency

```

n.initFingers(s1)
  f' = s1.f
  ∀ i ∈ 1..M s.th. (fini.start ∈ ]n, s1]),
    fini.node = s1
  ∀ j ∈ 1..M s.th. (finj.start ∉ ]n, s1]),
    finj.node = localSuccessor(f', finj.start)
n.localSuccessor(f, k)
  for i = 1..M
    if (k ∈ ]n, fini])
      return(fini)
  return(nil)
n.fixFingers(k)
  1 ≤ i = random() ≤ M
  fini.node =
    findSuccessor(fini.start)

```

Fig. 2. Initialization and Stabilization of Fingers

```

n.findSuccessor(k)
  //Case A: k is exactly equal to n
  if (k == n)
    return(n)
  //Case B: k is between n and s1
  if (k ∈ ]n, s1])
    return(firstAliveSuccessorNoChange());
  //Case C: Forward to the lookup to
  //the closest preceding alive finger
  cpf = closestAlivePrecedingFinger(k);
  if (cpf == nil)
    y = firstAliveSuccessorNoChange();
    if (k ∈ ]n, y])
      return(y);
  cpf = closestAlivePrecedingSucc(k);
  return(cpf.findSuccessor(k));
else
  return (cpf.findSuccessor(k));
n.firstAliveSuccessorNoChange()
  i = 1
  while (true)
    if (si == nil)
      //Broken Ring!!
    if (isAlive(si))
      return (si)
    i++
n.closestAlivePrecedingFinger(k)
  for i = M..1
    if ((fini ∈ ]n, k])
      and (fini ≠ nil)
      and isAlive(fini)
      return(fini)
  return(nil)
n.closestAlivePrecedingSucc(k)
  for i = S..1
    if ((si ∈ ]n, k])
      and (si ≠ nil)
      and isAlive(si)
      return(si)
  return(cpf)

```

Fig. 3. The Lookup Algorithm

are the main quantities of interest in the evaluation of any DHT. A node  $n$  looking up the successor of  $k$  runs the *findSuccessor* algorithm which can lead to the following cases:

*Case A.* If  $k$  is equal to  $n$  then  $n$  is trivially the successor of  $k$ .

*Case B.* If  $k \in ]n, s_1]$  then  $n$  has found the successor of  $k$ , but it could be that  $s_1$  has failed and  $n$  has not yet discovered this. However, entries in the successor list can act as backups for the first successor. Therefore, the first alive successor of  $n$  is the successor of  $k$ . Note that, in this case, while we try to find the first alive successor, we do not change the entries in the successor list. This is mainly because, to simplify the analysis, we want the successor list to be changed at a fixed rate rate  $\alpha\lambda_s$  only by the *fixSuccessors* function.

*Case C.* The lookup should be forwarded to a node closer to  $k$ , namely the closest alive finger preceding  $k$  in  $n$ 's finger table. The call to the function *closestAlivePrecedingFinger* returns such a node if possible and the lookup is forwarded to it. However, it could be the case that all alive preceding fingers



to  $k$  are dead. In that case, we need to use the successors list as a last resort for the lookup. Therefore, we locate the first alive successor  $y$  and if  $k \in ]n, y]$  then  $y$  is the successor of  $k$ . Otherwise, we locate the closest alive preceding successor to  $k$  and forward the lookup to it.

### C. Failures

Throughout the code we use the call *isAlive* and *isNotAlive*. A simple interpretation of those routines would be to equate them to a performance of a ping. However, a correct implementation for them is that they are discovered by performing the operation required. For instance, a call to *firstAliveSuccessor* in Fig. 1 is performed to retrieve a node  $y$  and then call *y.iThinkIamYourPred*, so alternatively the first alive successor could be discovered by iterating on the successor list and calling *iThinkIamYourPred*.

## IV. THE ANALYSIS

### A. Distributional Properties of Inter-Node Distances

During churn, the average inter-node distance is a fluctuating quantity whose distribution is used throughout our analysis. The derivation we present here of this distribution is independent of any details of the DHT implementation and depends solely on the dynamics of the join and leave process. It is hence applicable to any DHT that deploys a circular key space.

**Definition 4.1:** Given two keys  $u, v \in \{0 \dots \mathcal{K} - 1\}$ , the “distance” between them is  $u - v$  (with modulo- $\mathcal{K}$  arithmetic). We interchangeably say that  $u$  and  $v$  form an “interval” of length  $u - v$ . Hence the number of keys *inside* an interval of length  $\ell$  is  $\ell - 1$  keys.

**Definition 4.2:** Let  $Int_x$  be the number of intervals of length  $x$ , i.e. the number of pairs of consecutive nodes which are separated by a distance of  $x$  keys on the ring.

**Theorem 4.1:** For a process in which nodes join or leave with equal rates independently of each other and uniformly on the ring, and the number of nodes  $N$  in the network is almost constant with  $N \ll K$ , the probability ( $P(x) \equiv \frac{Int_x}{N}$ ) of finding an interval of length  $x$  is:  $P(x) = \rho^{x-1}(1 - \rho)$  where  $\rho = \frac{\mathcal{K} - N}{\mathcal{K}}$ .

*Proof:* By definition  $\sum P(x) = 1$  and  $\sum x P(x) = \mathcal{K}/N$ . Further, for the mean number of peers, the join-leave process we consider, simply implies that  $\frac{dN}{dt} = \lambda_j - \lambda_f$ . We will hence need to check that an equation for  $Int(x)$  does indeed satisfy all the above constraints. Note that the interval of time considered in the equation for the rate of change of  $N$  is the time-scale over which a *single* node change occurs. If we were to write the same equation for time-scales over which  $N$  node changes occur, the equation would then be  $\frac{dN}{dt} = (\lambda_j - \lambda_f)N$ . For our purposes however, we want to look at the changes in the system over “microscopic” time-scales in which at most one event occurs to change the state of the system.

We now write an equation for  $Int_x$  by considering all the processes which lead to its gain or loss. These are summarized in table I

First, a failure of either of the boundary nodes of an interval of size  $x$  leads to its loss at rate  $c_{1.1}$ . That is, since the node

$Int_x(t + \Delta t)$	Rate of Change
$= Int_x(t) - 1$	$c_{1.1} = (\lambda_f \Delta t) 2P(x)$
$= Int_x(t) - 1$	$c_{1.2} = (\lambda_j \Delta t) \frac{N(x-1)P(x)}{\mathcal{K}-N}$
$= Int_x(t) + 1$	$c_{1.3} = (\lambda_f \Delta t) \sum_{x_1=1}^{x-1} P(x_1)P(x-x_1)$
$= Int_x(t) + 1$	$c_{1.4} = (\lambda_j \Delta t) \frac{2N}{\mathcal{K}-N} \sum_{x_1 > x} P(x_1)$
$= Int_x(t)$	$1 - (c_{1.1} + c_{1.2} + c_{1.3} + c_{1.4})$

TABLE I  
GAIN AND LOSS TERMS FOR  $Int(x)$  THE NUMBER OF INTERVALS OF LENGTH  $x$ .

killed is randomly picked amongst all the nodes in the interval, the probability that it was participating on either side of an interval of length  $x$  is  $2P(x)$ .

Second, an interval of size  $x$  can be lost at rate  $c_{1.2}$  if a joining node splits it. Only joining with keys that belong to one of the  $Int_x$  intervals can lead to the loss of an interval of length  $x$  and in each one of these, there are  $x - 1$  ways (available keys) for splitting. Therefore  $(x-1) \times Int_x$  positions out of the  $\mathcal{K} - N$  available keys can destroy an interval of length  $x$ . That is, the probability that one of the intervals of length  $x$  is destroyed is  $\frac{(x-1)Int_x}{\mathcal{K}-N}$  which can be rewritten as  $\frac{N(x-1)P(x)}{\mathcal{K}-N}$ .

Third, the number of intervals of size  $x$  can increase by 1 at rate  $c_{1.3}$  if a failure of a boundary node results in the aggregation of two adjacent intervals. To clarify that, we give the following examples. An interval of length 1 cannot be formed by such a process. An interval of length 2 can be formed by the failure of a node if the node that failed was shared between two adjacent intervals of length 1. We are assuming here that the probability of picking two adjacent intervals of length 1 is  $P(1)P(1)$ . This is in effect assuming that the probability of having two adjacent intervals of size 1, factorises to  $P(1)^2$ . However for this system, this is an accurate estimation. Thus, in general, the probability of forming an interval of length  $x$  is  $\sum_{x_1=1}^{x-1} P(x_1)P(x-x_1)$ .

Fourth, an increase can happen at rate  $c_{1.4}$  if a join event splits a larger interval into an interval of size  $x$ . For a join to form an interval of length  $x$ , it must occur in an interval of length greater than  $x$ . In each interval of length  $x_1 > x$ , there are exactly two ways of forming an interval of length  $x$ . Therefore, the probability of forming an interval of length  $x$  is equal to  $\frac{2 \sum_{x_1 > x} Int_{x_1}}{\mathcal{K}-N}$ , which can be rewritten as  $\frac{2N \sum_{x_1 > x} P(x_1)}{\mathcal{K}-N}$ .

Finally,  $Int_x$  remains the same if none of the above happens.

Therefore the equation for  $Int_x$  for  $x > 1$  is:

$$\begin{aligned} \frac{dInt_x}{dt} = & -P(x) \left[ 2\lambda_f + \frac{N\lambda_j(x-1)}{\mathcal{K}-N} \right] \\ & + \lambda_f \sum_{x_1=1}^{x-1} P(x_1)P(x-x_1) \\ & + 2\lambda_j \frac{N}{\mathcal{K}-N} \sum_{x_1 > x} P(x_1). \end{aligned} \quad (1)$$

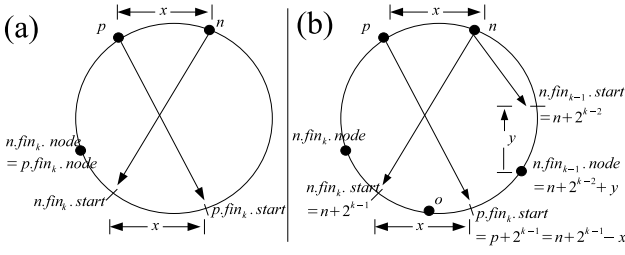


Fig. 4. (a) Case when  $n$  and  $p$  have the same value of  $fin_k.node$ . (b) Case where a newly joined node  $p$  copies the  $k$ -th entry of its successor node  $n$  as the best approximation for its own  $k$ -th entry (by the join protocol). In this case, there could be a node  $o$  which is the 'correct' entry for  $p.fin_k.node$ . However, since  $p$  is newly joined, the only information it has access to is the finger table of  $n$ .

The equation for  $Int_1$  is the same as the above except that the second term is missing.

We can check that :

$$\frac{d}{dt} \sum Int_x = \frac{dN}{dt} = \lambda_j - \lambda_f \quad (2)$$

as required.

Further we can check that the constraint:

$$\frac{d}{dt} \sum x Int_x = \frac{dK}{dt} = 0$$

is also obeyed. Equation 1 can be readily solved in the case  $\lambda_j = \lambda_f$  for the steady state (when the time derivative is zero) leading to the solution:

$$P(x) = \rho^{x-1}(1 - \rho) \quad (3)$$

where  $\rho = \frac{K-N}{K}$ .

Given the above term for  $\rho$  we can state the following corollary that gives an intuitive meaning for  $\rho$  in the case  $\lambda_j = \lambda_f$ .

**Corollary 1.1:** Given a ring of  $K$  keys populated by  $N$  nodes,  $\rho \equiv \frac{K-N}{K}$  is the ratio of the unpopulated keys to the total number of keys, i.e. the probability of picking a key at random and finding it empty is  $\rho$ .

The proof of the above theorem does assume that (in the case  $\lambda_j = \lambda_f$ ) the number of nodes  $N$  is fairly constant. Indeed at first sight this seems to be strictly true from Eq. 2. However, just as in a random walk, the variance in this case increases with time. We will comment more on the properties of the variance later. For the moment, we note that the above result can be generalized to also include the case when  $N$  is a fluctuating quantity. In this case we only need to multiply the  $N$  dependent terms in Eq. 1 with  $Prob(N, t)$ : the probability that there are  $N$  nodes in the system at time  $t$ , and average over  $N$ .

We now derive some properties of this distribution which will be used in the ensuing analysis.

**Property 4.1:** For any two keys  $u$  and  $v$ , where  $v = u + x$ , let  $b_i$  be the probability that the first node encountered in between these two keys is at  $u + i$  (where  $0 \leq i < x$ ). Then  $b_i \equiv \rho^i(1 - \rho)$ . The probability that there is definitely at least one node between  $u$  and  $v$  is:  $a(x) \equiv 1 - \rho^x$ . Hence the conditional probability that the first node is at a distance  $i$  given that there is at least one node in the interval is  $bc(i, x) \equiv b(i)/a(x)$ .

**Explanation :** Consider  $b_i$  first. For any key  $u$ , the probability that the first node encountered is at  $u$  itself ( $b_0$ ) is  $1 - \rho$  from Corollary 1.1. Similarly the probability that the first node encountered is at  $u + 1$  ( $b_1$ ) is  $\rho(1 - \rho)$  which is just the product of the probabilities that the first key is empty and the second is occupied. Thus in general, the probability that the first populated node starting from  $u$  is at  $u + i$  is  $b(i) \equiv (\rho)^i(1 - \rho)$ . Given this, the probability that there is at least one node between  $u$  and  $v = u + x$  (not including the case when the node is at  $v$ ) is  $\sum_{i=0}^{x-1} b_i = 1 - \rho^x \equiv a(x)$ . ■

**Property 4.2:** The probability that a node and at least one of its immediate predecessors share the same  $k$ -th finger is  $p_1(k) \equiv \frac{\rho}{1+\rho}(1 - \rho^{2^{k-2}})$ . This is  $\sim 1/2$  for  $K \gg 1$  and  $N \ll K$ . Clearly  $p_1 = 0$  for  $k = 1$ . It is straightforward (though tedious) to derive similar expressions for  $p_2(k)$  the probability that a node and atleast two of its immediate predecessors share the same  $k$ -th finger,  $p_3(k)$  and so on.

**Explanation :** If the distance between node  $n$  and its predecessor  $p$  is  $x$ , the distance between  $n.fin_k.start$  and  $p.fin_k.start$  is also  $x$  (see Fig. 4(a)). If there is no node in between  $n.fin_k.start$  and  $p.fin_k.start$  then  $n.fin_k.node$  and  $p.fin_k.node$  will share the same value. From Eq. 3, the probability that the distance between  $n$  and  $p$  is  $x$  is  $\rho^{x-1}(1 - \rho)$ . However,  $x$  has to be less than  $2^{k-1}$ , otherwise  $p.fin_k.node$  will be equal to  $n$ . The probability that no node exists between  $n.fin_k.start$  and  $p.fin_k.start$  is  $\rho^x$  (by Property 4.1). Therefore the probability that the  $n.fin_k.node$  and  $p.fin_k.node$  share the same value is:  $\sum_{x=1}^{2^{k-1}-1} \rho^{x-1}(1 - \rho)\rho^x = \frac{\rho}{1+\rho}(1 - \rho^{2^{k-2}})$  ■

**Property 4.3:** We can similarly assess the probability that the join protocol (Section refsec:fingers) results in further replication of the  $k$ -th pointer. Let us define the probability  $p_{join}(i, k)$  as the probability that a newly joined node, chooses the  $i$ -th entry of its successor's finger table for its own  $k$ -th entry. Note that this is unambiguous even in the case that the successor's  $i$ -th entry is repeated. All we are asking is, when is the  $k$ -th entry of the new joinee the same as the  $i$ -th entry of the successor? Clearly  $i \leq k$ . Infact for the larger fingers, we need only consider  $p_{join}(k, k)$ , since  $p_{join}(i, k) \sim 0$  for  $i < k$ . Using the interval distribution we find, for large  $k$ ,  $p_{join}(k, k) \sim \rho(1 - \rho^{2^{k-2}-2}) + (1 - \rho)(1 - \rho^{2^{k-2}-2}) - (1 - \rho)\rho(2^{k-2} - 2)\rho^{2^{k-2}-3}$ . This function goes to 1 for large  $k$ .

**Explanation :** By the join protocol a newly joined node  $p$ , tries to assign  $p.fin_k.node$  to the best approximate value from the finger table of its successor  $n$ . This approximate value might turn out to be  $n.fin_k.node$ , especially for the larger fingers. If  $p$  chooses the  $k$ -th entry of  $n$  as its own  $k$ -th entry, it must be because the  $k - 1$ -th entry of  $n$  (if distinct, as is always the case for large  $k$ ) does not afford it a better choice. The condition for this is:  $p.fin_k.start > n.fin_{k-1}.node$ . If the distance between  $n.fin_k.start$  and  $p.fin_k.start$  is  $x$ , and the distance between  $n.fin_{k-1}.start$  and  $n.fin_{k-1}.node$  is  $y$  (see Fig. 4 (b)), then the constraint on  $x$  and  $y$  is  $n + 2^{k-2} + y > n + 2^{k-2} + x$  or  $x + y < 2^{k-2}$ . We also have the added constraint that  $x < 2^{k-1}$ , since otherwise  $p.fin_k.node$  would simply be

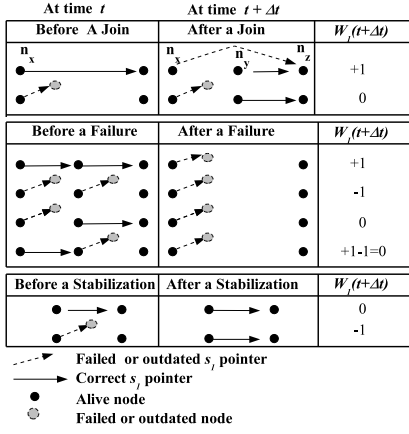


Fig. 5. Changes in  $W_1$ , the number of wrong (failed or outdated)  $s_1$  pointers, due to joins, failures and stabilizations.

$n$ . In fact since the distance between the  $n_{\text{fin}_k.\text{start}}$  and  $n_{\text{fin}_{k-1}.\text{start}}$  cannot be more than  $2^{k-2}$  we have  $x < 2^{k-2}$ .

Thus the probability  $p_{\text{join}}(k, k)$  is:

$$\sum_{x=1}^{2^{k-2}-1} \sum_{y=1}^{2^{k-2}-x} P(x)P(y) = \sum_{z=2}^{2^{k-2}-1} \rho^{z-2}(1-\rho)^2(z-1) \quad (4)$$

where we have put in the expressions for  $P(x)$  and  $P(y)$  from Eq. 3 and converted the double summation to a single one. This expression can be summed easily to obtain the result quoted above.

We can also analogously compute  $p_{\text{join}}(i, k)$  for any  $i$ . The only trick here is to estimate the probability that starting from  $i$ , the last *distinct* entry of  $n$ 's finger table *does not* give  $p$  a better choice for its  $k_{\text{th}}$  entry. This can again readily be computed using property 4.1.

### B. Successor Pointers

We now turn to estimating various quantities of interest for Chord. In all that follows we will evaluate various *average* quantities, as a function of the parameters. However this same formalism can also be used for evaluating higher moments like the variance.

In the case of Chord, we need consider only one of three kinds of events happening at any micro-instant: a join, a failure or a stabilization. One assumption made in the following is that such a micro-instant of time exists, or in other words, that we can divide time till we have an interval small enough that in this interval, only any one of these three processes occur. Implicit in this is the assumption that a stabilization (either of successors or fingers) is over much faster than the time-scales over which joins and fails occur. Another (more serious) assumption is that the state of the system is a *product* of the state of all the nodes. Nodes are hence assumed to have, for the most part, states independent of each other, *i.e.* the probability of two adjacent nodes having a wrong successor pointer is taken to be the product of the individual nodes having wrong successor pointers (though as we have seen from Properties 4.2 and 4.3, in the case of finger pointers, we

Change in $W_1(r, \alpha)$	Rate of Change
$W_1(t + \Delta t) = W_1(t) + 1$	$c_{2.1} = (\lambda_j \Delta t)(1 - w_1)$
$W_1(t + \Delta t) = W_1(t) + 1$	$c_{2.2} = \lambda_f(1 - w_1)^2 \Delta t$
$W_1(t + \Delta t) = W_1(t) - 1$	$c_{2.3} = \lambda_f w_1^2 \Delta t$
$W_1(t + \Delta t) = W_1(t) - 1$	$c_{2.4} = \alpha \lambda_s w_1 \Delta t$
$W_1(t + \Delta t) = W_1(t)$	$1 - (c_{2.1} + c_{2.2} + c_{2.3} + c_{2.4})$

TABLE II

GAIN AND LOSS TERMS FOR  $W_1(r, \alpha)$ : THE NUMBER OF WRONG FIRST SUCCESSORS AS A FUNCTION OF  $r$  AND  $\alpha$ .

do also consider the case when adjacent nodes might have correlated fingers). These assumptions imply that the analysis is not *exact*. However as we see below it is sufficiently precise to predict all quantities extremely accurately.

Consider first the successor pointers. Let  $w_k(r, \alpha)$ ,  $d_k(r, \alpha)$  denote the fraction of nodes having a *wrong*  $k^{\text{th}}$  successor pointer or a *failed* one respectively and  $W_k(r, \alpha)$ ,  $D_k(r, \alpha)$  be the respective *numbers*. A *failed* pointer is one which points to a departed node and a *wrong* pointer points either to an incorrect node (alive but not correct) or a dead one. As we will see, both these quantities play a role in predicting lookup consistency and lookup length.

By the protocol for stabilizing successors in Chord, a node periodically contacts its first successor, possibly correcting it and reconciling with its successor list. Therefore, the number of wrong  $k^{\text{th}}$  successor pointers are not independent quantities but depend on the number of wrong first successor pointers. We first consider  $s_1$  here, and then briefly discuss the other cases towards the end of this section.

We write an equation for  $W_1(r, \alpha)$  by accounting for all the events that can change it in a micro event of time  $\Delta t$ . An illustration of the different cases in which changes in  $W_1$  take place due to joins, failures and stabilizations is provided in Fig. 5. In some cases  $W_1$  increases/decreases while in others it stays unchanged. For each increase/decrease, Table II provides the corresponding probability.

By our implementation of the join protocol, a new node  $n_y$ , joining between two nodes  $n_x$  and  $n_z$ , has its  $s_1$  pointer always correct after the join. However the state of  $n_x.s_1$  before the join makes a difference. If  $n_x.s_1$  was correct (pointing to  $n_z$ ) before the join, then after the join it will be wrong and therefore  $W_1$  increases by 1. If  $n_x.s_1$  was wrong before the join, then it will remain wrong after the join and  $W_1$  is unaffected. Thus, we need to account for the former case only. The probability that  $n_x.s_1$  is correct is  $1 - w_1$  and from that follows the term  $c_{2.1}$ .

For failures, we have 4 cases. To illustrate them we use nodes  $n_x$ ,  $n_y$ ,  $n_z$  and assume that  $n_y$  is going to fail. First, if both  $n_x.s_1$  and  $n_y.s_1$  were correct, then the failure of  $n_y$  will make  $n_x.s_1$  wrong and hence  $W_1$  increases by 1. Second, if  $n_x.s_1$  and  $n_y.s_1$  were both wrong, then the failure of  $n_y$  will decrease  $W_1$  by one, since one wrong pointer disappears. Third, if  $n_x.s_1$  was wrong and  $n_y.s_1$  was correct, then  $W_1$  is unaffected. Fourth, if  $n_x.s_1$  was correct and  $n_y.s_1$  was wrong, then the wrong pointer of  $n_y$  disappears and  $n_x.s_1$  becomes wrong, therefore  $W_1$  is unaffected. For the first case to happen, we need to pick two nodes with correct pointers,

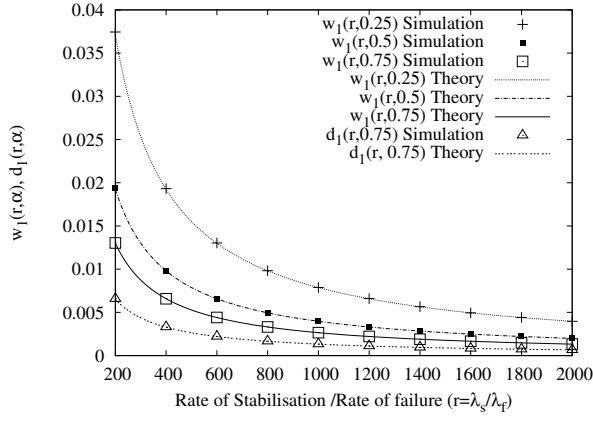


Fig. 6. Theory and Simulation for  $w_1(r, \alpha)$ ,  $d_1(r, \alpha)$

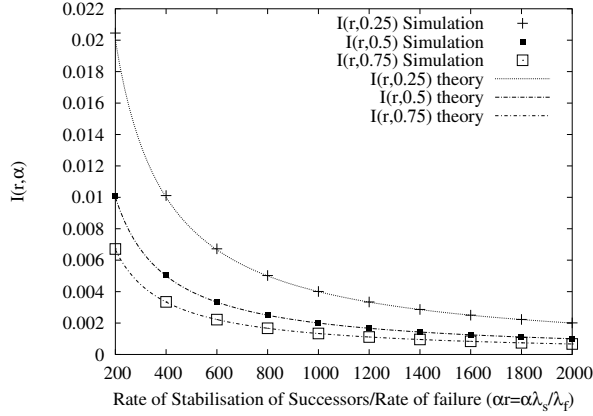


Fig. 7. Theory and Simulation for  $I(r, \alpha)$

the probability of this is  $(1 - w_1)^2$ . For the second case to happen, we need to pick two nodes with wrong pointers, the probability of this is  $w_1^2$ . From these probabilities follow the terms  $c_{2.2}$  and  $c_{2.3}$ .

Finally, a successor stabilization does not affect  $W_1$ , unless the stabilizing node had a wrong pointer. The probability of picking such a node is  $w_1$ . From this follows the term  $c_{2.4}$ .

Hence the equation for  $W_1(r, \alpha)$  is:

$$\frac{dW_1}{dt} = \lambda_j(1 - w_1) + \lambda_f(1 - w_1)^2 - \lambda_f w_1^2 - \alpha \lambda_s w_1$$

Solving for  $w_1$  in the steady state and putting  $\lambda_j = \lambda_f$ , we get:

$$w_1(r, \alpha) = \frac{2}{3 + r\alpha} \approx \frac{2}{r\alpha} \quad (5)$$

This expression matches well with the simulation results as shown in Fig. 6.  $d_1(r, \alpha)$  is then  $\approx \frac{1}{2}w_1(r, \alpha)$  since when  $\lambda_j = \lambda_f$ , about half the number of wrong pointers are incorrect and about half point to dead nodes. Thus  $d_1(r, \alpha) \approx \frac{1}{r\alpha}$  which also matches well the simulations as shown in Fig. 6. We can also use the above reasoning to iteratively get  $w_k(r, \alpha)$  for any  $k$ .

### C. Break-up (Network Disconnection) Probability

We demonstrate below, how calculating  $d_k(r, \alpha)$ : the fraction of nodes with dead  $k^{th}$  pointers, helps in estimating

Change in $W_1(r, \alpha)$	Rate of Change
$N_{bu}(t + \Delta t) = N_{bu}(t) + 1$	$c_{3.1} = (\lambda_f \Delta t) d_1(r, \alpha)$
$N_{bu}(t + \Delta t) = N_{bu}(t) + 1$	$c_{3.2} = \lambda_f \Delta t (1 - d_1) d_2$
$N_{bu}(t + \Delta t) = N_{bu}(t) - 1$	$c_{3.3} = \alpha \lambda_s \Delta t P_{bu}(2, r, \alpha)$
$N_{bu}(t + \Delta t) = N_{bu}(t)$	$1 - (c_{3.1} + c_{3.2} + c_{3.3})$

TABLE III

GAIN AND LOSS TERMS FOR  $N_{bu}(2, r, \alpha)$ : THE NUMBER OF NODES WITH DEAD FIRST AND SECOND SUCCESSORS

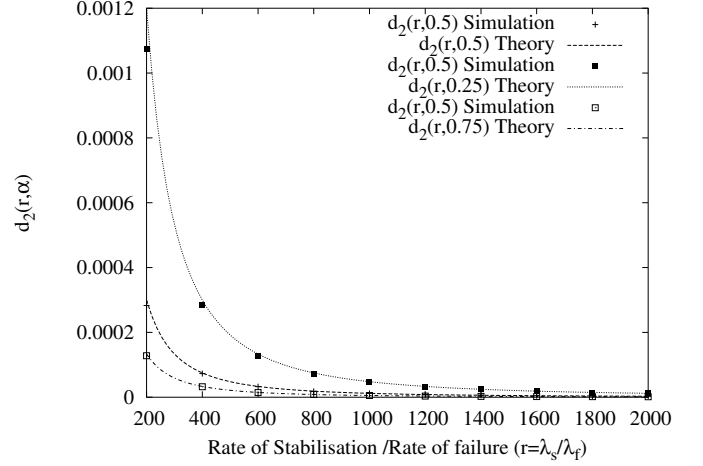


Fig. 8. Theory and Simulation for  $d_2(r, \alpha)$

precisely the probability that the network gets disconnected for any value of  $r$  and  $\alpha$ . Let  $P_{bu}(n, r, \alpha)$  be the probability that  $n$  consecutive nodes fail. If  $n = \mathcal{S}$ , the length of the successor list, then clearly the node whose successor list this is gets disconnected from the network and the network breaks up. For the range of  $r$  considered in Fig. 6,  $P_{bu}(\mathcal{S}, r, \alpha) \sim 0$ . However should we go lower, this starts becoming finite. The master equation analysis introduced here can be used to estimate  $P_{bu}(n, r, \alpha)$  for any  $1 \leq n \leq \mathcal{S}$ . We indicate how this might be done by considering the case  $n = 2$ . Let  $N_{bu}(2, r, \alpha)$  be the number of configurations in which a node has both  $s_1$  and  $s_2$  dead and  $P_{bu}(2, r, \alpha)$  be the fraction of such configurations. Table III indicates how this is estimated within the present framework.

A join event does not affect this probability in any way. So we need only consider the effect of failures or stabilization events. The term  $c_{3.1}$  accounts for the situation when the *first* successor of a node is dead (which happens with probability  $d_1(r, \alpha)$  as explained above). A failure event can then kill its second successor as well and this happens with probability  $c_{3.1}$ . The second term is the situation that the first successor is alive (with probability  $1 - d_1$ ) but the second successor is dead (with probability  $d_2$ ). This probability is  $\sim 2/\alpha r$ . (the second successor of a node being dead either implies that the first successor of *its* first successor is dead with probability  $d_1$ , or that it has not stabilized recently, and hence has not corrected its second successor pointer. This happens with probability  $\sim 1/\alpha r$ . These two terms add up to  $2/\alpha r$ ). A stabilization event reduces the number of such configurations by one, if the node doing the stabilization had such a configuration to begin with.



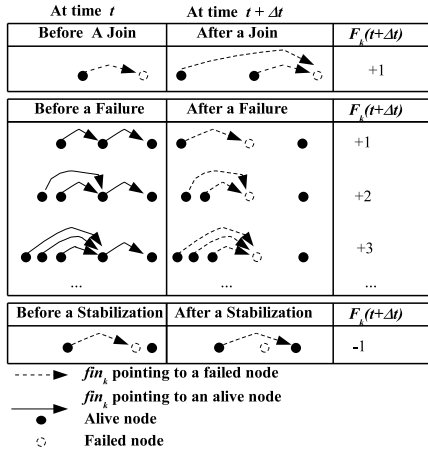


Fig. 9. Changes in  $F_k$ , the number of failed  $fin_k$  pointers, due to joins, failures and stabilizations.

Solving the equation for  $N_{bu}(2, r, \alpha)$ , one hence obtains that  $P_{bu}(2, r, \alpha) \sim 3/(\alpha r)^2$ . As Fig. 8 shows, this is a precise estimate.

We can similarly estimate the probabilities for three consecutive nodes failing, *etc.*, and hence also the disconnection probability  $P_{bu}(\mathcal{S}, r, \alpha)$ . This formalism thus affords the possibility of making a precise prediction for when the system runs the danger of getting disconnected as a function of the parameters.

**Lookup Consistency** By the lookup protocol, a lookup is inconsistent if the immediate predecessor of the sought key has a wrong  $s_1$  pointer. However, we need only consider the case when the  $s_1$  pointer is pointing to an alive (but incorrect) node since our implementation of the protocol always requires the lookup to return an alive node as an answer to the query. The probability that a lookup is inconsistent  $I(r, \alpha)$  is hence  $w_1(r, \alpha) - d_1(r, \alpha)$ . This prediction matches the simulation results very well, as shown in Fig. 7.

#### D. Failure of Fingers

We now turn to estimating the fraction of finger pointers which point to failed nodes. As we will see this is an important quantity for predicting lookups, since failed fingers cause timeouts and increase the lookup length. We need however only consider fingers pointing to *dead* nodes. Unlike members of the successor list, *alive* fingers even if outdated, always bring a query closer to the destination and do not affect consistency or substantially even the lookup length. Therefore we consider fingers in only two states, alive or dead (failed). By our implementation of the stabilization protocol (see Sections III-A and III-B), fingers and successors are stabilized entirely independently of each other to simplify the analysis. Thus even though the first finger is also always the first successor, this information is not used by the node in updating the finger.

Let  $f_k(r, \alpha)$  denote the fraction of nodes having their  $k^{th}$  finger pointing to a failed node and  $F_k(r, \alpha)$  denote the respective number. For notational simplicity, we write these as simply  $F_k$  and  $f_k$ . We can predict this function for any  $k$  by again estimating the gain and loss terms for this quantity,

$F_k(t + \Delta t)$	Rate of Change
$= F_k(t) + 1$	$c_{4.1} = (\lambda_j \Delta t) \sum_{i=1}^k p_{join}(i, k) f_i$
$= F_k(t) - 1$	$c_{4.2} = (1 - \alpha) \frac{1}{\mathcal{M}} f_k (\lambda_s \Delta t)$
$= F_k(t) + 1$	$c_{4.3} = (1 - f_k)^2 [1 - p_1(k)] (\lambda_f \Delta t)$
$= F_k(t) + 2$	$c_{4.4} = (1 - f_k)^2 (p_1(k) - p_2(k)) (\lambda_f \Delta t)$
$= F_k(t) + 3$	$c_{4.5} = (1 - f_k)^2 (p_2(k) - p_3(k)) (\lambda_f \Delta t)$
$= F_k(t)$	$1 - (c_{4.1} + c_{4.2} + c_{4.3} + c_{4.4} + c_{4.5})$

TABLE IV

SOME OF THE RELEVANT GAIN AND LOSS TERMS FOR  $F_k$ , THE NUMBER OF NODES WHOSE  $k^{th}$  FINGERS ARE POINTING TO A FAILED NODE FOR  $k > 1$ .

caused by a join, failure or stabilization event, and keeping only the most relevant terms. These are listed in table IV and illustrated in Fig. 9

A join event can play a role here by increasing the number of  $F_k$  pointers if the successor of the joiner had a failed  $i^{th}$  pointer (occurs with probability  $f_i$ ) and the joiner replicated this from the successor as the joiner's  $k^{th}$  pointer. (occurs with probability  $p_{join}(i, k)$  from property 4.3). For large enough  $k$ , this probability is one only for  $p_{join}(k, k)$ , that is the new joiner mostly only replicates the successor's  $k^{th}$  pointer as its own  $k^{th}$  pointer. This is what we consider here.

A stabilization evicts a failed pointer if there was one to begin with. The stabilization rate is divided by  $\mathcal{M}$ , since a node stabilizes any one finger randomly, every time it decides to stabilize a finger at rate  $(1 - \alpha)\lambda_s$ .

Given a node  $n$  with an alive  $k^{th}$  finger (occurs with probability  $1 - f_k$ ), when the node pointed to by that finger fails, the number of failed  $k^{th}$  fingers ( $F_k$ ) increases. The amount of this increase depends on the number of immediate predecessors of  $n$  that were pointing to the failed node with their  $k^{th}$  finger. That number of predecessors could be 0, 1, 2,... etc. Using property 4.2 the respective probabilities of those cases are:  $1 - p_1(k)$ ,  $p_1(k) - p_2(k)$ ,  $p_2(k) - p_3(k)$ ,... etc.

Solving for  $f_k$  in the steady state, we get:

$$f_k = \frac{\left[ 2\tilde{P}_{rep}(k) + 2 - p_{join}(k) + \frac{r(1-\alpha)}{\mathcal{M}} \right]}{2(1 + \tilde{P}_{rep}(k))} - \frac{\sqrt{\left[ 2\tilde{P}_{rep}(k) + 2 - p_{join}(k) + \frac{r(1-\alpha)}{\mathcal{M}} \right]^2 - 4(1 + \tilde{P}_{rep}(k))^2}}{2(1 + \tilde{P}_{rep}(k))} \quad (6)$$

where  $\tilde{P}_{rep}(k) = \sum p_i(k)$ . In principle its enough to keep even three terms in the sum. The above expressions match very well with the simulation results (Fig. 11).

#### E. Cost of Finger Stabilizations and Lookups

In this section, we demonstrate how the information about the failed fingers and successors can be used to predict the cost of stabilizations, lookups or in general the cost for reaching any key in the id space. By cost we mean the number of hops needed to reach the destination *including* the number of timeouts encountered en-route. Timeouts occur every time a query is passed to a dead node. The node does not answer and the originator of the query has to use another finger instead.



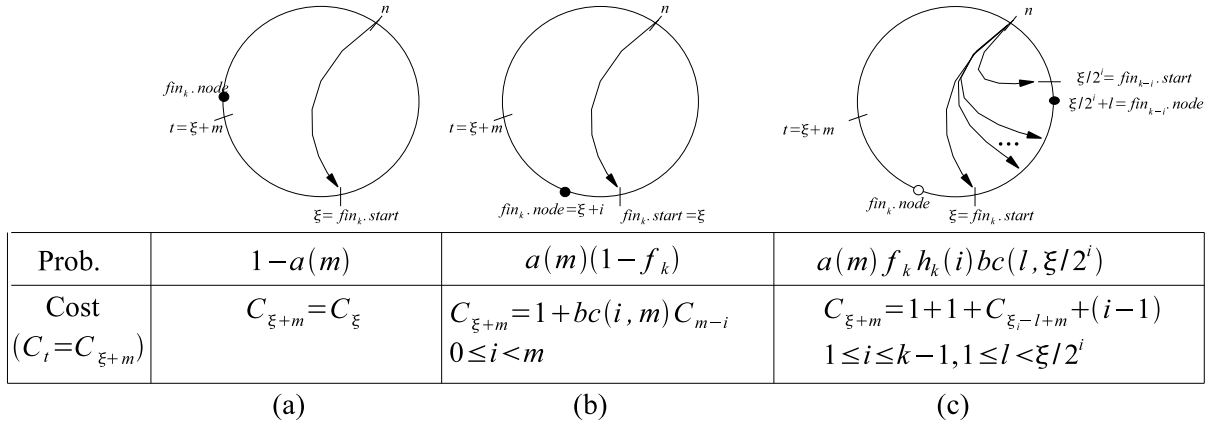


Fig. 10. Cases that a lookup can encounter with the respective probabilities and costs.

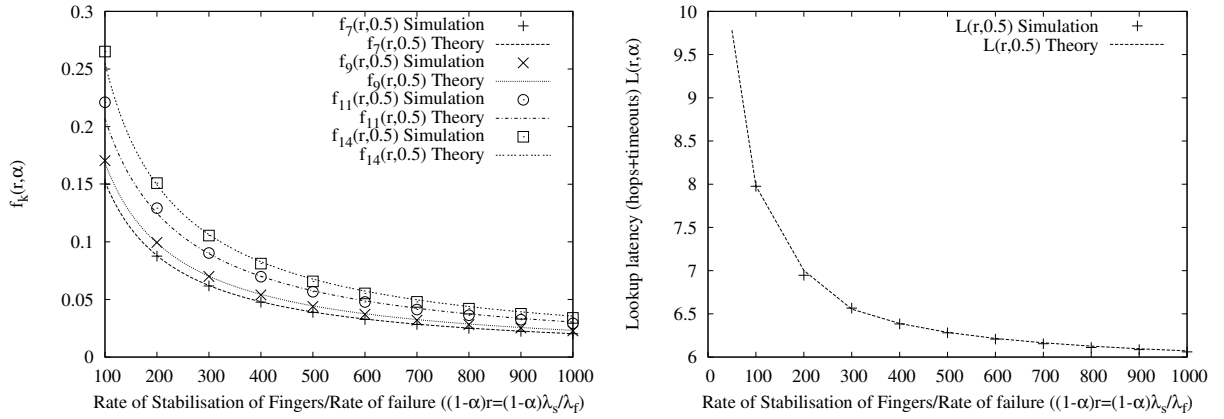


Fig. 11. Theory and Simulation for  $f_k(r, \alpha)$ , and  $L(r, \alpha)$

For this analysis, we consider timeouts and hops to add equally to the cost. We can easily generalize this analysis to investigate the case when a timeout costs some factor  $\gamma$  times the cost of a hop.

Define  $C_t(r, \alpha)$  (also denoted  $C_t$ ) to be the expected cost for a given node to reach some target key which is  $t$  keys away from it (which means reaching the first successor of this key). For example,  $C_1$  would then be the cost of looking up the adjacent key (1 key away). Since the adjacent key is always stored at the first alive successor, therefore if the first successor is alive (which occurs with probability  $1 - d_1$ ), the cost will be 1 hop. If the first successor is dead but the second is alive (occurs with probability  $d_1(1 - d_2)$ ), the cost will be 1 hop + 1 timeout = 2 and the *expected* cost is  $2 \times d_1(1 - d_2)$  and so forth. Therefore, we have  $C_1 = 1 - d_1 + 2 \times d_1(1 - d_2) + 3 \times d_1d_2(1 - d_3) + \dots \approx 1 + d_1 = 1 + 1/(\alpha r)$ .

For finding the expected cost of reaching a general distance  $t$  we need to follow closely the Chord protocol, which would lookup  $t$  by first finding the closest preceding finger. For the purposes of the analysis, we will find it easier to think in terms of the closest preceding *start*. Let us hence define  $\xi$  to be the start of the finger (say the  $k^{th}$ ) that most closely precedes  $t$ . Hence  $\xi = 2^{k-1} + n$  and  $t = \xi + m$ , i.e. there are  $m$  keys between the sought target  $t$  and the start of the most closely preceding finger. With that, we can write a recursion relation

for  $C_{\xi+m}$  as follows:

$$\begin{aligned}
 C_{\xi+m} &= C_{\xi} [1 - a(m)] \\
 &+ (1 - f_k)a(m) \left[ 1 + \sum_{i=0}^{m-1} bc(i, m)C_{m-i} \right] \\
 &+ f_k a(m) \left[ 1 + \sum_{i=1}^{k-1} h_k(i) \right. \\
 &\quad \left. \sum_{l=0}^{\xi/2^i-1} bc(l, \xi/2^i)(1 + (i-1) + C_{\xi-l+m}) + O(h_k(k)) \right]
 \end{aligned} \tag{7}$$

where  $\xi_i \equiv \sum_{m=1, i} \xi/2^m$  and  $h_k(i)$  is the probability that a node is forced to use its  $k - i^{th}$  finger owing to the death of its  $k^{th}$  finger. The probabilities  $a, b, bc$  have already been introduced in Section IV, and we define the probability  $h_k(i)$  below.

The lookup equation though rather complicated at first sight merely accounts for all the possibilities that a Chord lookup will encounter, and deals with them exactly as the protocol dictates.

The first term (Figure 10 (a)) accounts for the eventuality that there is no node intervening between  $\xi$  and  $\xi + m$  (occurs with probability  $1 - a(m)$ ). In this case, the cost of looking for  $\xi + m$  is the same as the cost for looking for  $\xi$ .

The second term (Figure 10 (b)) accounts for the situation when a node does intervene inbetween (with probability  $a(m)$ ), and this node is alive (with probability  $1 - f_k$ ). Then the query is passed on to this node (with 1 added to register the increase in the number of hops) and then the cost depends on the length of the distance between this node and  $t$ .

The third term (Figure 10 (c)) accounts for the case when the intervening node is dead (with probability  $f_k$ ). Then the cost increases by 1 (for a timeout) and the query needs to find an alternative lower finger that most closely precedes the target. Let the  $k - i^{th}$  finger (for some  $i$ ,  $1 \leq i \leq k - 1$ ) be such a finger. This happens with probability  $h_k(i)$ , i.e., the probability that the lookup is passed back to the  $k - i^{th}$  finger either because the intervening fingers are dead or share the same finger table entry as the  $k^{th}$  finger is denoted by  $h_k(i)$ . The start of the  $k - i^{th}$  finger is at  $\xi/2^i$  and the distance between  $\xi/2^i$  and  $\xi$  is equal to  $\sum_{m=1,i} \xi/2^m$  which we denote by  $\xi_i$ . Therefore, the distance from the start of the  $k - i^{th}$  to the target is equal to  $\xi_i + m$ . However, note that  $fin_{k-i}.node$  could be  $l$  keys away (with probability  $bc(l, \xi/2^i)$ ) from  $fin_{k-i}.start$  (for some  $l$ ,  $0 \leq l < \xi/2^i$ ). Therefore, after making one hop to  $fin_{k-i}.node$ , the remaining distance to the target is  $\xi_i + m - l$ . The increase in cost for this operation is  $1 + (i - 1)$ ; the 1 indicates the cost of taking up the query again by  $fin_{k-i}.node$ , and the  $i - 1$  indicates the cost for trying and discarding each of the  $i - 1$  intervening fingers. The probability  $h_k(i)$  is easy to compute given property 4.1 and the expression for the  $f_k$ 's computed in the previous section.

$$\begin{aligned} h_k(i) &= a(\xi/2^i)(1 - f_{k-i}) \\ &\quad \times \prod_{s=1,i-1} (1 - a(\xi/2^s) + a(\xi/2^s)f_{k-s}), i < k \quad (8) \\ h_k(k) &= \prod_{s=1,k-1} (1 - a(\xi/2^s) + a(\xi/2^s)f_{k-s}) \end{aligned}$$

Equation .8 accounts for all the reasons that a node may have to use its  $k - i^{th}$  finger instead of its  $k^{th}$  finger. This could happen because the intervening fingers were either dead or not distinct. The probabilities  $h_k(i)$  satisfy the constraint  $\sum_{i=1}^k h_k(i) = 1$  since clearly, either a node uses any one of its fingers or it doesn't. This latter probability is  $h_k(k)$ , that is the probability that a node cannot use any earlier entry in its finger table. In this case,  $n$  proceeds to its successor list. The query is now passed on to the first alive successor and the new cost is a function of the distance of this node from the target  $t$ . We indicate this case by the last term in equation 7 which is  $O(h_k(k))$ . This can again be computed from the inter-node distribution and from the functions  $d_k(r, \alpha)$  computed earlier. However in practice, the probability for this is extremely small except for targets very close to  $n$ . Hence this does not significantly affect the value of general lookups and we ignore it for the moment.

The cost for general lookups is hence

$$L(r, \alpha) = \frac{\sum_{i=1}^{\mathcal{K}-1} C_i(r, \alpha)}{\mathcal{K}}$$

The lookup equation is solved recursively numerically, given the coefficients and  $C_1$ . We plot the result in Fig 11. The theoretical result matches the simulation very well.

## F. Analysis of the Lookup Equation in the zero-churn case

On general grounds, it is easy to argue that the average lookup cost has the following form  $A + \frac{B}{r} + \frac{C}{r^2} + \dots$ . The dependence on churn is specified by the  $r$ -dependence and  $A, B$  etc depend on the other parameters of the system like  $N$  and  $\mathcal{K}$ . To get  $A$ , we need to consider equation 7 with no churn (all  $f_k$ 's set to zero). To get  $B$ , we need to analyze the lookup equation to  $O(\frac{1}{r})$  and so on. In the following section, we study the lookup equation 7 in some detail to understand the behaviour without churn. This is useful in order to ascertain that it does indeed reproduce known results such as for example, that the average lookup cost is  $0.5 * \log(N)$  without churn [10]. Infact as we will see, for any  $N$ , the average lookup cost as predicted by equation 7 is indeed  $0.5 * \log(N)$  plus some  $\rho$ -dependent corrections which though small are accurately predicted. An added benefit of the analysis is that we can also predict what the average lookup without churn will be for any base (Chord has base 2 and accordingly has a finger table size of  $\log_2(\mathcal{K})$ ). By our definition of higher bases a system of base  $b$  will have a finger table size of  $(b - 1)\log_b(\mathcal{K})$ .

Equation 7 with the churn-dependent terms set to zero becomes:

$$C_{\xi+m} = C_{\xi} [1 - a(m)] + a(m) + \sum_{i=0}^{m-1} b(i)C_{m-i} \quad (9)$$

After some rewriting of this, it is easily seen that the cost for any key  $i + 1$  can be written as the following recursion relation:

$$C_{i+1} = \rho C_i + (1 - \rho) + (1 - \rho)C_{i+1-\xi(i+1)} \quad (10)$$

Here we have used the definition of  $a$  and  $b$  from the internode-interval distribution and the notation  $\xi(i + 1)$  refers to the start of the finger most closely preceding  $i + 1$ . For instance, for  $i + 1 = 4$ ,  $\xi(i + 1) = 2$  and for  $i + 1 = 11$ ,  $\xi(i + 1) = 8$  etc.

In figure 12, we have plotted  $C_i$  versus  $i$  by solving equation 10 numerically.

We are interested in solving the recursion relation and computing  $L = \frac{1}{\mathcal{K}} \sum_{i=1}^{\mathcal{K}-1} C_i$ . To do this, we decompose this sum into the following partial sums:

$$\begin{aligned} s_0 &= C_1 = 1 \\ s_1 &= C_2 \\ s_2 &= C_3 + C_4 \\ s_3 &= C_5 + C_6 + C_7 + C_8 \\ &\dots \\ s_{\mathcal{M}} &= C_{2^{\mathcal{M}-1}+1} + \dots + C_{\mathcal{K}-1} \end{aligned} \quad (11)$$

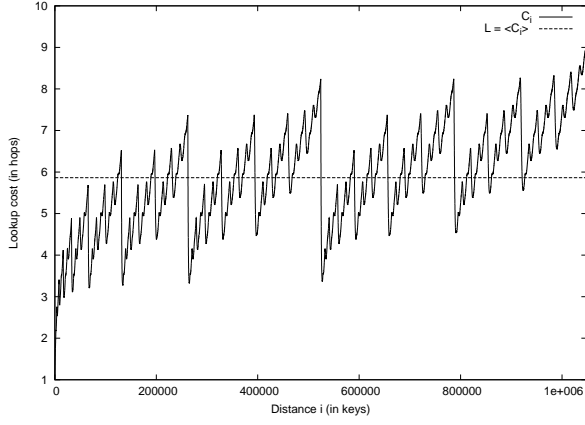


Fig. 12. The average cost  $C_i$  (the number hops for looking up an item  $i$  keys away) in a network of  $N = 1000$  nodes and  $K = 2^{20}$  keys without churn obtained from the recurrence relation (10). The average lookup length  $L$  is also plotted as a reference.

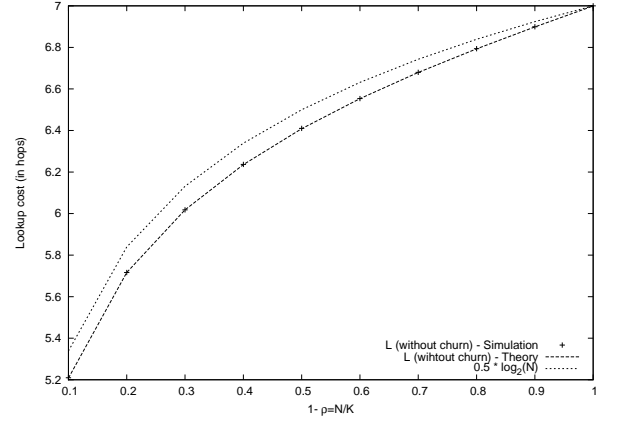


Fig. 13. Theory and Simulation for the lookup cost without churn for a key space of size  $K = 2^{14}$  for varying  $N$ . Plotted as reference is the curve  $0.5 \log_2(N)$ . Note that on the y axis we have actually plotted  $L - 1$  for convenience.

Substituting the expressions for the  $C$ 's in the above, we find:

$$\begin{aligned}
 s_0 &= 1 \\
 s_1 &= \frac{\rho}{1-\rho} [C_1 - C_2] + 1 + s_0 \\
 s_2 &= \frac{\rho}{1-\rho} [C_2 - C_4] + 2 + [s_0 + s_1] \\
 &\dots \\
 s_i &= \frac{\rho}{1-\rho} [C_{2^{i-1}} - C_{2^i}] + 2^{i-1} + \sum_{j=0}^{i-1} s_j
 \end{aligned} \quad (12)$$

By substituting serially the expressions for  $s_j$  (where  $0 \leq j \leq i-1$ ), the expression for  $s_i$  (for  $i \geq 2$ ) becomes:

$$\begin{aligned}
 s_i &= \frac{\rho}{1-\rho} [2^{i-2} C_1 - C_{2^i} - \sum_{j=1}^{i-2} s^{i-2-j} C_{2^j}] \\
 &\quad + 2^i + (i-1)2^{i-2}
 \end{aligned} \quad (13)$$

Hence

$$\begin{aligned}
 \sum_{i=0}^M s_i &= -\rho + [2^{M+1} - 1] + M2^{M-1} - [2^M - 1] \\
 &\quad + \frac{\rho}{1-\rho} \left[ (2^{M-1} - 1)C_1 - \sum_{i=2}^{M-1} C_{2^i} - C_{K-1} \right. \\
 &\quad \left. - (2^{M-2} - 1)C_2 - (2^{M-3} - 1)C_4 - \dots \right]
 \end{aligned} \quad (14)$$

Therefore

$$\begin{aligned}
 \sum_{i=0}^M s_i &= -\rho + 2^M + M2^{M-1} \\
 &\quad + \frac{\rho}{1-\rho} \left[ (2^{M-1} - 1)C_1 - \sum_{i=2}^{M-1} C_{2^i} - C_{K-1} \right. \\
 &\quad \left. - \sum_{j=2}^{M-2} (2^{M-j} - 1)C_{2^{j-1}} \right]
 \end{aligned} \quad (15)$$

The equation for the average lookup length without churn is thus,

$$\begin{aligned}
 L &= \frac{\sum s}{K} \\
 &= -\frac{\rho}{K} + 1 + \frac{1}{2}M \\
 &\quad + \frac{\rho}{1-\rho} \left[ \frac{2^{M-1} - 1}{K} C_1 - \frac{1}{K} \sum_{i=2}^{M-1} C_{2^i} - \frac{1}{K} C_{K-1} \right. \\
 &\quad \left. - \sum_{j=2}^{M-2} \frac{2^{M-j} - 1}{K} C_{2^{j-1}} \right]
 \end{aligned} \quad (16)$$

If we can take the limit  $K \rightarrow \infty$ , we can throw away some of the terms.

$$\begin{aligned}
 \lim_{K \rightarrow \infty} L &= 1 + \frac{1}{2}M \\
 &\quad + \frac{\rho}{1-\rho} \left[ \frac{C_1}{2} - \frac{1}{K} \sum_{i=1}^{M-1} C_{2^i} + \frac{C_2}{K} - \frac{1}{K} C_{K-1} \right. \\
 &\quad \left. - \sum_{j=2}^{M-2} \frac{2^{M-j}}{K} C_{2^{j-1}} + \sum_{j=2}^{M-2} \frac{C_{2^{j-1}}}{K} \right] \\
 &\approx 1 + \frac{1}{2}M + \frac{\rho}{1-\rho} \left[ \frac{C_1}{2} - \frac{C_2}{4} - \frac{C_4}{8} \dots - \frac{C_{2^{M-3}}}{2^{M-2}} \right]
 \end{aligned} \quad (17)$$

Since  $C_1 = 1$ , we can write

$$\begin{aligned}
 L &= 1 + \frac{1}{2}M - \frac{\rho}{2(1-\rho)} \left[ \frac{C_2 - 1}{2} + \frac{C_4 - 1}{4} + \dots \right. \\
 &\quad \left. + \frac{C_{2^{M-3}} - 1}{2^{M-3}} \right]
 \end{aligned} \quad (18)$$

From the recursion relation for the  $C_i$ 's, it is easy to see that

$$(C_i - 1) = (1-\rho)g_i^{(1)}(\rho) + (1-\rho)^2 g_i^{(2)}(\rho) + \dots \quad (19)$$

where the  $g_i$ 's are functions only of  $\rho$ .

Hence if  $(1-\rho)$  is small ( $\frac{N}{K} \rightarrow 0$ ), we need only compute the  $C_i$ 's to first order in  $(1-\rho)$  to get the leading order effect and second order in  $(1-\rho)$  to get the correction etc.

Hence in general the, the expression for  $L$  is:

$$L = 1 + \frac{1}{2}\mathcal{M} - \frac{\rho}{2} \left[ e_1(\rho) + (1-\rho)e_2(\rho) + (1-\rho)^2e_3(\rho) \dots \right] \quad (20)$$

Where  $e_1(\rho) = \sum_{i=1}^{\mathcal{M}-3} g_{2^i}^{(1)}(\rho)$  etc.

We evaluate this expression numerically by solving recursion relation (10) and compare it with simulations done at zero churn. As can be seen the prediction of the equation is very accurate (Figure 13).

Let us now compute  $e_1(\rho)$  to see what the leading order effect is. We now need to solve recursion relation (10) only to order  $1-\rho$ , which gives:

$$\begin{aligned} C_2 - 1 &= (1-\rho) \\ C_4 - 1 &= (1-\rho) [1 + \rho + \rho^2] \\ C_8 - 1 &= (1-\rho) [1 + \rho + \rho^2 + \dots + \rho^6] \\ &\dots \\ C_i - 1 &= (1-\rho) [1 + \rho + \rho^2 + \dots + \rho^{i-2}] \end{aligned} \quad (21)$$

Therefore,

$$L = 1 + \frac{1}{2}\mathcal{M} + \frac{\rho}{2} \left[ \frac{1}{2} + \frac{1+\rho+\rho^2}{4} + \dots \right] \quad (22)$$

Consider the expression inside the brackets. We are computing this in the approximation  $\frac{N}{K} = \epsilon \rightarrow 0$ , i.e.  $\rho = 1-\epsilon$ , therefore  $\rho^x = (1-\epsilon)^x \approx e^{-\epsilon x}$ . If  $x > \frac{1}{\epsilon}$ , then  $\rho^x \rightarrow 0$ , therefore if  $x > \frac{K}{N}$ , then  $\rho^x \rightarrow 0$ . Hence, the terms inside the brackets become:

$$\sum_{j=1}^T \frac{2^j - 1}{2^j} + (2^T - 1) \sum_{j=T+1}^{\mathcal{M}-3} \frac{1}{2^j} \quad (23)$$

Where  $T \equiv \ln_2 K - \ln_2 N$  and we have put  $\rho^x \approx 1$  for  $x < \frac{K}{N}$  and  $\rho \rightarrow 0$  for  $x > \frac{K}{N}$ . This is clearly an overestimation and so we expect the result to over estimate the exact expression 20.

Expression 23 becomes:

$$T - \left[ 1 - \left( \frac{1}{2} \right)^{\mathcal{M}-3} \right] + \left[ 1 - \left( \frac{1}{2} \right)^{\mathcal{M}-3-T} \right] \approx T$$

Therefore:

$$\begin{aligned} L &= 1 + \frac{1}{2} \ln_2 K - \frac{1}{2} [\ln_2 K - \ln_2 N] \\ &\approx 1 + \frac{1}{2} \ln_2 N \end{aligned} \quad (24)$$

Which is the known result for the average lookup length of Chord.

Another important parameter in the performance of DHTs in general is the base. By increasing the base, the number of fingers per node increases which leads to a shorter lookup path length. The effect of varying the base has been studied in [2], [6]. So far, we have considered in this analysis base-2 Chord. We can likewise carry out this analysis for any base.

In general, we have base- $b$  with  $(b-1)\log_b(K)$  fingers per node. Consider as an example  $b = 4$ . Here we can define the

the partial sums again in the following manner:

$$\begin{aligned} \Delta_0 &= s_0 = C_1 = 1 \\ \Delta_1 &= s_1 + s_2 + s_3 \\ \Delta_2 &= s_4 + s_5 + s_6 \\ &\dots \end{aligned} \quad (25)$$

where

$$\begin{aligned} s_1 &= C_2 = \rho C_1 + (1-\rho) + (1-\rho)C_1 \\ s_2 &= C_3 = \rho C_2 + (1-\rho) + (1-\rho)C_1 \\ s_3 &= C_4 = \rho C_3 + (1-\rho) + (1-\rho)C_1 \\ s_4 &= C_5 + C_6 + C_7 + C_8 \\ s_5 &= C_9 + C_{10} + C_{11} + C_{12} \\ s_6 &= C_{13} + C_{14} + C_{15} + C_{16} \\ &\dots \end{aligned} \quad (26)$$

Therefore

$$\begin{aligned} \Delta_0 &= C_1 \\ \Delta_1 &= \rho [\Delta_1 + C_1 - C_4] + 3(1-\rho) + 3(1-\rho) [\Delta_0] \\ \Delta_2 &= \rho [\Delta_2 + C_4 - C_{16}] + 12(1-\rho) + 3(1-\rho) [\Delta_0 + \Delta_1] \\ &\dots \end{aligned} \quad (27)$$

In general for a base  $b$ , define  $B \equiv b-1$  and  $b^{\mathcal{M}} = K$ . Then we have:

$$\begin{aligned} \Delta_j &= \frac{\rho}{1-\rho} [C_{b^{j-1}} - C_{b^j}] \\ &+ B(B+1)^{j-1} + B [\Delta_0 + \Delta_1 + \dots + \Delta_{j-1}] \end{aligned} \quad (28)$$

Following much the same procedure as before, we find

$$\begin{aligned} L &= \frac{1}{K} \sum_{j=0}^{\mathcal{M}} \Delta_j \\ &\approx 1 + \frac{B}{B+1} \mathcal{M} - \frac{B}{B+1} \frac{\rho}{1-\rho} \left[ \frac{C_b - 1}{B+1} + \frac{C_{b^2} - 1}{(B+1)^2} + \dots \right] \end{aligned} \quad (29)$$

for  $K \rightarrow \infty$  as the analogue of (18). Again we can simplify and slightly overestimate the sum by assuming that  $\rho^x \approx 0$  for  $x > \frac{K}{N}$  and  $\rho^x \approx 1$  for  $x < \frac{K}{N}$ . Then we get:

$$L \approx 1 + \frac{b-1}{b} \frac{\ln_2 N}{\ln_2 b} \quad (30)$$

This is the analogue of equation 24 for any base  $b$ .

Clearly it is of interest to carry out a similar analysis with churn to get an estimate of the  $O(1/r)$  effect. However in this case there is no simple analogue of equation 10. The principle complication comes from the last term in equation 7 the 'back-tracking' term which accounts for a node not using the closest preceding finger to the target, owing to its failure, but an earlier one. This results in the recursion relation for  $C(i+1)$  depending on not just two earlier costs (costs to reach two keys closer to the node in question that  $i+1$ ) as in equation 10 but on a larger and larger number of earlier terms as  $i$  increases. We are nevertheless investigating this further.



## V. WHAT IS CHURN?

We now discuss a broader issue, connected with churn, which arises naturally in the context of our analysis. As we mentioned earlier, all our analysis is performed in the steady state where the rate of joins ( $\lambda_j$ ) is equal to the rate of failures  $\lambda_f$ . However the rates  $\lambda_j$  and  $\lambda_f$  can themselves each be chosen in one of two different ways. They could either be “per-network” or “per-node”. In the former case, the number of joiners (or the number of failures) *does not* depend on the current number of nodes in the network. This is the case when a poisson model is considered either for arrivals or departures. Put in another way, this is like saying that on average, there is always a fixed number of nodes joining or failing per time interval, irrespective of the total number of nodes in the network. In the case when these rates are chosen to be per-node, the number of joiners or failures *does* depend on the current number of occupied nodes). We consider three possibilities here, when  $\lambda_j$  is per-network and  $\lambda_f$  is per-node; both are per-network or (as is the case studied in this paper) both are per-node. In all three cases, since the system is always studied in the steady state where the total number of joiners per unit time is equal to the total number of failures per unit time, the equation for the mean is always  $dN/dt = 0$ . We hence expect the mean behavior to be the same, at least in the regime when  $N$  is roughly constant. However the behavior of fluctuations is very different in each of these three cases. As mentioned earlier, the time-scale over which the rate of change of  $N$  is evaluated is again a ‘microscopic’ time scale with a single node change occurring at every interval of time.

In the first case, the steady state condition is  $\lambda_j/N_o = \lambda_f$ , where  $N_o$  is the initial number of nodes in the system. The equation for the mean is  $dN/dt = \lambda_j/N - \lambda_f$ , which ensures that  $N$  cannot deviate too much from the steady state value. Similarly one can write an equation for the second moment  $N^2$ :  $dN^2/dt = (\lambda_j/N + \lambda_f) + 2(\lambda_j - N\lambda_f)$ . While the first term is a ‘noise’ term which encourages fluctuations, the second term becomes stronger the larger the deviation from  $N_o$  and hence strongly damps out fluctuations. Thus the number of nodes in the system remains close to its initial value.

In the second case, where the join and failure rates are both per-network the equation for the mean is  $dN/dt = \lambda_j/N - \lambda_f/N$ . Hence putting  $\lambda_j = \lambda_f$  ensures the steady state condition. However in this case, the equation for the second moment is  $dN^2/dt = (\lambda_j/N + \lambda_f/N)$ . The joins-failures process thus makes the system execute a “random-walk” in  $N$ , where the “steps” of the walk depend on  $N$  and are smaller if  $N$  is larger. For such a system, fluctuations are not bounded and a large deviation can and will take the system to the  $N = 0$  state eventually. The time for this to happen scales with  $N$  as  $N^3$  for this process.

The third case (which is also the case considered in this paper) is when both rates are per-node. This is very similar to the second case. The equation for the mean is just  $dN/dt = \lambda_j - \lambda_f$  as mentioned earlier. Again setting  $\lambda_j = \lambda_f$  ensures steady state. The equation for the second moment is now  $dN^2/dt = (\lambda_j + \lambda_f)$ . There is thus again no “repair” mechanism for large fluctuations, and the system will be

eventually driven to extinction. In this case the process on  $N$  is just an ordinary random walk and the time taken to hit the  $N = 0$  state scales as  $N^2$ .

Which of these ‘types’ of churn is the most relevant? In the real world, the churn felt by a DHT, might possibly be some time-varying mixture of these three, and will also possibly depend on the application. It is hence probably of importance to study all these mechanisms and their implications in detail.

## VI. DISCUSSION AND CONCLUSION

To summarize, in this paper, we have presented a detailed theoretical analysis of a DHT-based P2P system, Chord, using a Master-equation formalism. This analysis differs from existing theoretical work done on DHTs in that it aims not at establishing bounds, but on precise determination of the relevant quantities in this dynamically evolving system. From the match of our theory and the simulations, it can be seen that we can predict with an accuracy of greater than 1% in most cases.

Though this analysis is not *exact* (in the sense that there are approximations made to make the analysis simpler), yet it provides a methodology to keep track of most of the relevant details of the system. We expect that the same analysis can be done for most other DHT’s in a similar manner, thus helping to establish quantitative guidelines for their comparison.

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